

FIG. 3

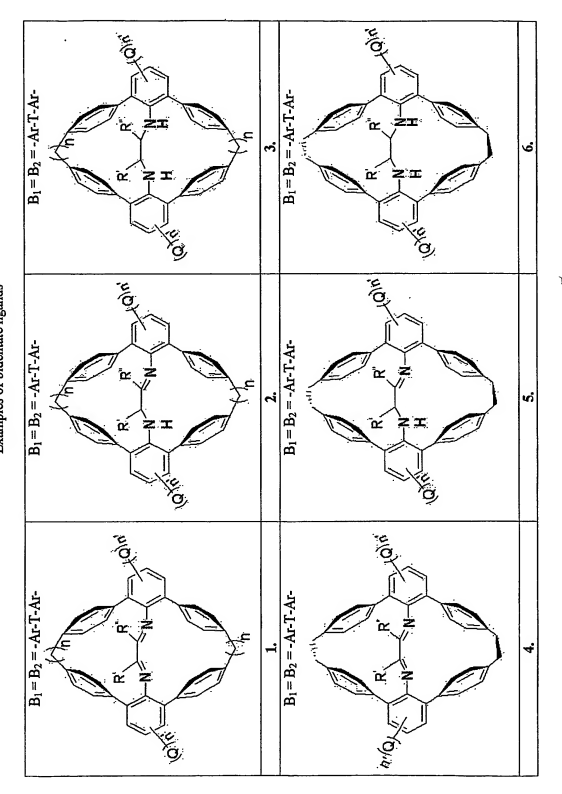
Summary of Polymerization Data

Branches Per 1000 carbons 1.45 1.43 1.41 386 619 342 323 429 294 307 286 333 263 331 Yield (g) 6.10 7.35 9.25 3.20 6.85 9.00 3.11 Time (min) Moles of Catalyst (x 10°) Entry

^aExperimental condition: in 200 mL of toluene, cocatalyst MMAO (Al:Ni ≈ 3000), 200 psi ethylene pressure.

^b TON = turnover number, which was calculated as the moles of ethylene per mole of catalyst; TOF = turnover frequency, i.e., TON per hour.

FIG. 4A
Examples of bidentate ligands



5/16

FIG. 4B Examples of bidentate ligands (continued)

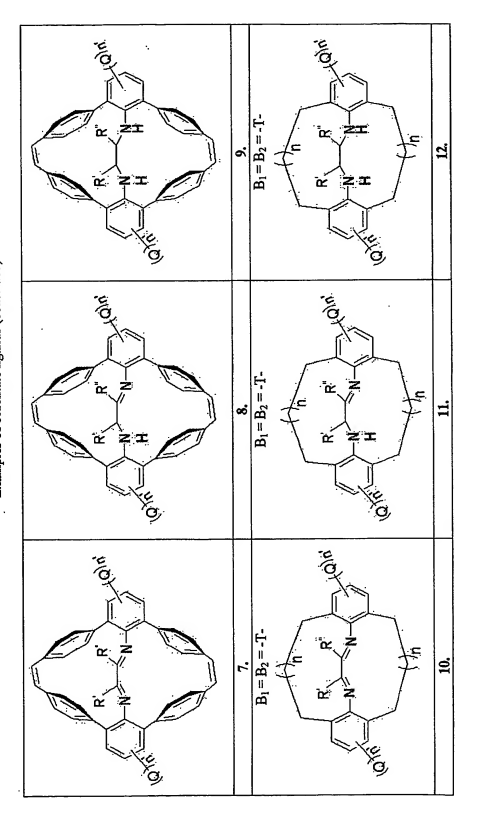
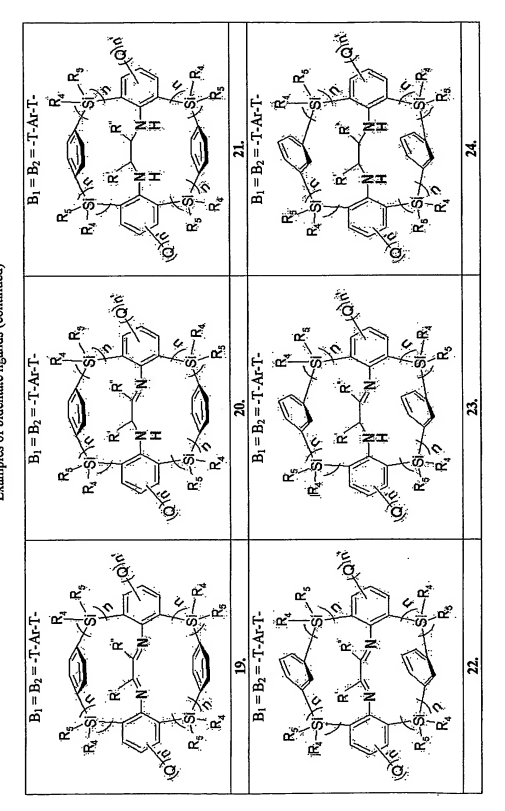


FIG. 4CExamples of bidentate ligands (continued)

| B ₁ = B ₂ = -T-Ar-T- R R R R H H H H H H H H H H H H H H H | 15. | B ₁ = B ₂ | 18. |
|---|-----|--|-----|
| $B_1 = B_2 = -T - Ar - T -$ R R N N N N | 14. | $B_1 = B_2 = -T \cdot Ar \cdot T \cdot$ $A = A \cdot Ar \cdot T \cdot$ $A \cdot A \cdot$ | 17. |
| B ₁ = B ₂ = -T-Ar-T- R R' R'' Ola R R R'' N AN R'' | 13, | $B_1 = B_2 = -T - Ar - T -$ R | 16. |

7/16

FIG. 4DExamples of bidentate ligands (continued)



half-cyclic structure for any of the above ligands $B_1 = B_2 = -T - Ar - T -$ FIG. 4E
Examples of bidentate ligands (continued) 26. 28. т Г $B_1 = B_2 = -T - Ar - T$ $B_1 = B_2 = -T - Ar - T -$ 25. 27.

9/16

FIG. 5A Examples of tridentate ligands

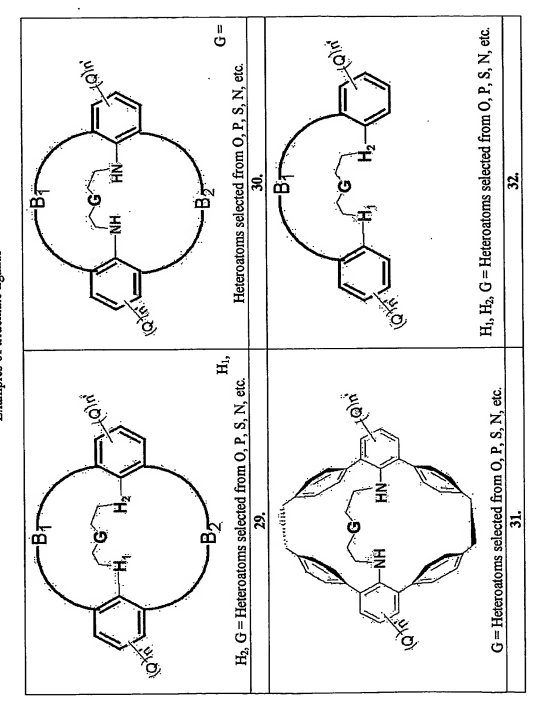


FIG. 5B
Examples of tridentate ligands (continued)

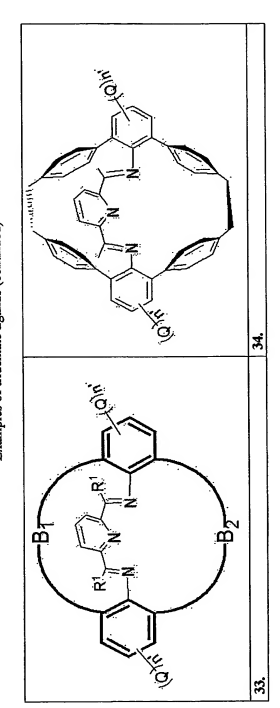


FIG. 6A

Examples of preference of metals for different types of ligands

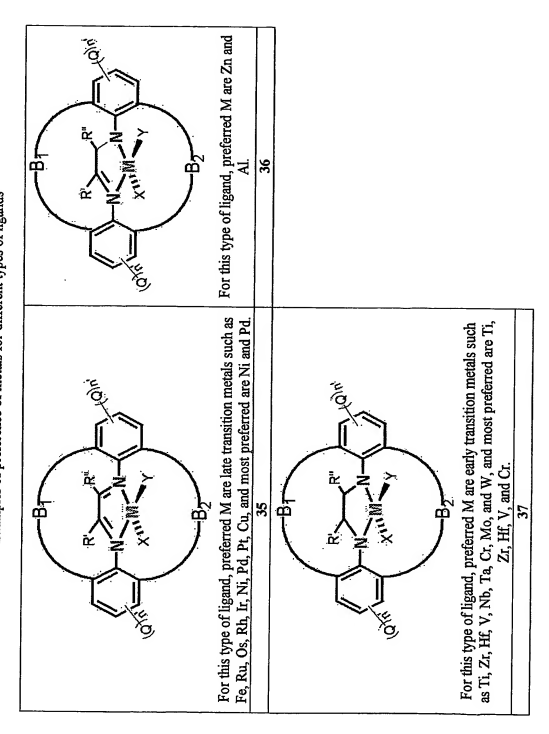


FIG. 6B

Examples of preference of metals for different types of ligands (continued)

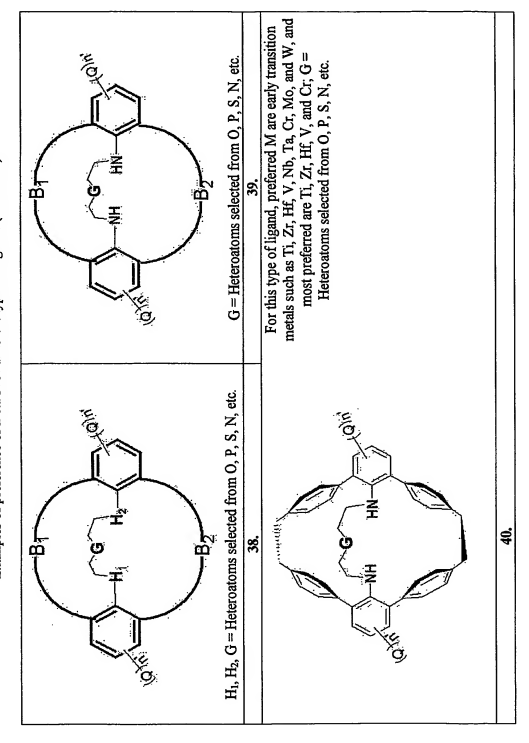
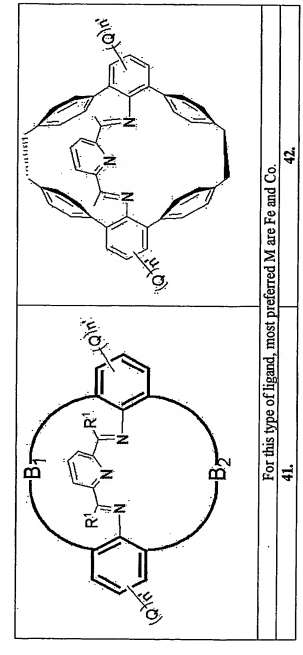


FIG. 6C

Examples of preference of metals for different types of ligands (continued)



In structures 1 through 42, Q, n, R4 and R5 are as defined in Formula 1 in the specification, n' is 1 through 4, and R' and R" are alkyl, alkenyl, aryl, aralkyl, or cycloalkyl.

WO 2005/014658 PCT/US2004/025586

14/16

FIG. 7
(Prior Art)

FIG. 8

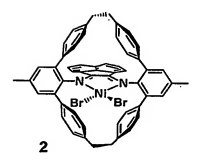
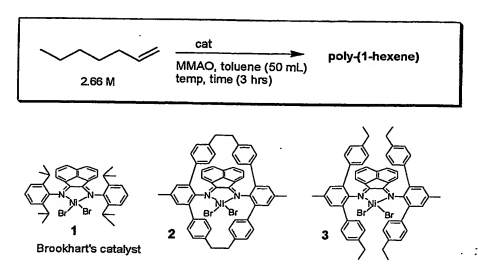


FIG. 9



15/16

FIG. 10

Polymerization of 1-Hexene
(Catalyst Activity and Molecular Weight Data at 0 °C and at 25 °C)

| entry | c at | Load (mmol) | Temp (oC) | Yield (g) | TON | Mw | Mn | PDI | DSC (°C) | Branch /1000c |
|---------------------------------|---------|--|--------------|--------------|------|-------|-------|------|--|------------------|
| F62 | 2 | 0.005 | 0 | 0.33 | 784 | 170 K | 96 K | 1.77 | T _m 58 | 65 |
| F36 | 2 | 0.01 | 0 | 0.41 | 488 | 305 K | 292 K | 1.05 | T _m 63 | 61 |
| F84 | 1 | 0.005 | 0 | 1.62 | 3850 | 719 K | 627 K | 1.15 | T _m -42 | 104 |
| Ref JACS 95, 6414 | 1 | 0.017 activated by Et ₂ AICI | 0 | 2.1 | 1468 | 310 K | 140 K | 2.2 | T _m -20 T _g -57 | 100 |
| F60 | 2 | 0.005 | 25 | 1.68 | 3992 | 623 K | 510 K | 1.22 | T _m 62 | 57 |
| F72 | 1 | 0.005 | 25 | 1.90 | 4515 | 817 K | 543 K | 1.50 | T _m -50 | 108 |
| Ref <i>JACS 96,11664</i> | 1 | 0.017 3.2 M; 30 min rxn. | 23 | | 2800 | 129 K | 84 K | 1.54 | T _m -17 T _g -57 | 120 |
| F74 | 3 | 0.005 | 25 | 0.80 | 1901 | 88 K | 83 K | 1.06 | T _m 56 | 35 |
| F42 | 2 | 0.005 | 75 | 2.21 | 5466 | 622 K | 529 K | 1.17 | T _m 59 | 52 |
| F68 | 1 | 0.005 | 75 | 0.43 | 1022 | 415 K | 279 K | 1.49 | T _m -53 | 111 |
| F70 | 3 | 0.005 | 75 | 0.26 | 618 | 131 K | 92 K | 1.43 | T _m 73 | 38 |
| F46 | 2 | 0.005 | 95 | 1.6 | 3802 | 252 K | 125 K | 2.00 | T _m 57 | 54 |
| F48 | 1 | 0.005 | 95 | 0.47 | 1117 | 287 K | 171 K | 1.68 | T _m -53 | 113 |
| F50 | 3 | 0.005 | 95 | 0.59 | 1402 | 77 K | 59 K | 1.29 | T _m 76 | 40 |

FIG. 11



